

# Pivotal Role of the Non-*hr* Origin of DNA Replication in the Genesis of Defective Interfering Baculoviruses

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**The generation of deletion mutants, including defective interfering viruses, upon serial passage of *Spodoptera exigua* multicapsid nucleopolyhedrovirus (SeMNPV) in insect cell culture has been studied. Sequences containing the non-homologous region origin of DNA replication (non-*hr ori*) became hypermolar in intracellular viral DNA within 10 passages in Se301 insect cells, concurrent with a dramatic drop in budded virus and polyhedron production. These predominant non-*hr ori*-containing sequences accumulated in larger concatenated forms and were generated de novo as demonstrated by their appearance and accumulation upon infection with a genetically homogenous bacterial clone of SeMNPV (bacmid). Sequences were identified at the junctions of the non-*hr ori* units within the concatemers, which may be potentially involved in recombination events. Deletion of the SeMNPV non-*hr ori* using RecE/RecT-mediated homologous ET recombination in *Escherichia coli* resulted in a recombinant bacmid with strongly enhanced stability of virus and polyhedron production upon serial passage in insect cells. This suggests that the accumulation of non-*hr oris* upon passage is due to the replication advantage of these sequences. The non-*hr ori* deletion mutant SeMNPV bacmid can be exploited as a stable eukaryotic heterologous protein expression vector in insect cells.**

Baculoviruses are large enveloped, circular double-stranded DNA insect viruses which are widely used as bioinsecticides in agriculture and forestry and can be genetically engineered to improve their effectiveness (2, 18). More recently, baculoviruses were shown to have potential as gene delivery vectors for gene therapy (12, 32, 45) or as vectors for surface display of complex eukaryotic proteins (6). Yet, their major application to date is as a viral vector for the expression of heterologous proteins in insect cells (19, 36). The prototypic and most intensively studied baculovirus, *Autographa californica* multicapsid nucleopolyhedrovirus (AcMNPV), has been primarily used as an expression vector, while other baculoviruses may become exploited as well, especially when appropriate cell lines are available.

A major drawback in the large-scale production of baculoviruses as bioinsecticides or for heterologous protein production is the so-called passage effect. This effect is notable as a significant drop in production after prolonged virus passaging in insect cell culture (reviewed by Krell [25]) and is a result of the accumulation of defective interfering particles (DIs) (20). These DIs are rapidly generated in cell culture (39) and become predominant after prolonged passaging, meanwhile interfering with the replication of intact helper virus. However, the mechanism of the generation of DIs is still enigmatic, and the sequences involved are unknown.

DIs have retained *cis*-acting elements essential for baculovirus DNA synthesis, such as origins of replication (*ori*) (25). Transient virus-mediated plasmid replication assays demon-

strated that baculovirus homologous regions (*hrs*) (21, 22, 28, 37), as well as baculovirus early promoter regions not containing *hr* sequences (49), have a putative *ori* function. In addition, these assays showed that other non-*hrs*, with structural similarities to eukaryotic *oris*, may have an *ori* function (11, 14, 23, 38). Their *ori* activity in vivo was recently demonstrated but was unfortunately not compared to the *ori* activity of *hrs* (7). Strikingly, AcMNPV DIs were enriched in such a non-*hr ori* (26, 27). This suggests a prominent role of baculovirus non-*hr oris* in the generation of DIs.

For large-scale or continuous production of heterologous eukaryotic proteins by use of the baculovirus expression system in insect cell bioreactors, the passage effect is a major obstacle. For prevention of the negative consequences of the passage effect, a genetically stable viral genotype is highly demanded. This may be achieved by selection for viruses with enhanced stability or higher polyhedron (or recombinant protein) production (42) or, alternatively, by site-directed mutagenesis of viral sequences putatively involved in the generation and/or maintenance of DIs. Therefore, we chose the non-*hr ori* sequence as a target for mutagenesis studies.

Compared to AcMNPV infections in widely used cell lines such as Sf21 and Sf9, the generation and predominance of *Spodoptera exigua* MNPV (SeMNPV) deletion mutants (including DIs) in various *S. exigua* cell lines occurs significantly faster (3, 11). This virus-cell system thus provides a better model system than AcMNPV to study the passage effect, the mechanism of DI generation, and the pivotal role of non-*hr ori* sequences therein.

In this paper we have studied the rapid passage effect during serial passaging of wild-type SeMNPV in the established *S. exigua* cell line Se301 (9). To monitor the generation of DI genomes over passage and to study the role of the non-*hr ori* sequences in this process, a full-length infectious clone of

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SeMNPV propagated in *Escherichia coli* (bacmid) was constructed and used in serial passage experiments. This revealed the pivotal role of the non-*hr ori* in the genesis of DIs and led to the generation of a recombinant SeMNPV bacmid with enhanced stability in cultured insect cells.

## MATERIALS AND METHODS

**Cells, insects, and virus.** The *S. exigua* cell line Se301 (8, 9) was donated by T. Kawarabata (Institute of Biological Control, Kyushu University, Kyushu, Japan) and was propagated at 27°C in Grace's supplemented medium (Gibco BRL) containing 10% fetal calf serum (Gibco BRL). Fourth-instar *S. exigua* larvae were infected by contamination of artificial diet with  $4 \times 10^5$  SeMNPV-US1 (5) polyhedra per larva (43). Hemolymph was collected as previously described (17) and was defined as the passage zero (P0) budded-virus (BV) inoculum to initiate serial passage in cultured Se301 cells. Serial undiluted passaging was carried out as previously described (39). Infectious BV titers were determined using the endpoint dilution assay (47).

**DNA isolation, Southern hybridization, colony lift, molecular cloning, and sequencing.** Intracellular viral (ICV) DNA and BV DNA were isolated as previously described (44). Digested viral DNA was run overnight in ethidium bromide-stained 0.6% agarose gels, and Southern blotting was performed by standard capillary upward blotting (40) using Hybond-N (Amersham Pharmacia) filters. As a DNA size marker,  $\lambda$ -DNA digested with *EcoRI/HindIII/BamHI* was used. Randomly primed DNA probes for Southern hybridization were made using a digoxigenin nonradioactive nucleic acid labeling and detection system (Roche). PCR products (927 bp) of the SeMNPV non-*hr ori* were made with reverse primer DZ127, 5'-CATCGATGCGTACGTGACTTTC-3' (nucleotides [nt] 84027 to 84048 [16]), and forward primer DZ128, 5'-CCTTGCCTTCCTTGGTG-3' (nt 83122 to 83139); purified using a High Pure PCR purification kit (Roche); and digoxigenin labeled overnight. Hybridization and colorimetric detection with nitroblue tetrazolium-5-bromo-4-chloro-3-indolylphosphate (Gibco BRL) were performed according to the manufacturer's recommendations. Hypermolar viral *XbaI* bands were cut from the gel, purified with Glassmax (Gibco BRL), and cloned into pUC19 by electroporation of *E. coli* DH5 $\alpha$  using standard methods (40). A colony lift assay (40) was used to isolate the cloned submolar 5.3-kb fragment using the same probe as described above. Automatic sequencing was performed using an ABI prism 310 genetic analyzer (Perkin-Elmer) at the Laboratory of Molecular Biology, Wageningen University. Sequence analyses were performed using BLAST (1) from the University of Wisconsin Genetics Computer Group computer programs (release 10.0).

**Construction of bacmid cloning vector.** The bacmid vector for direct cloning of SeMNPV was constructed by PCR using the Expand long-template PCR system (Roche). Custom made primers (Gibco BRL) were designed using DNASTar Primerselect and were based on the sequence of AcMNPV transfer plasmid pVL1393 (29), which was the backbone of the transfer vector pMON14272 used to construct the AcMNPV bacmid bMON14272 (30). Primers DZ113 (5'-CCTTCTGAGGTACCTTCTAGAAATCCGGAG-3') and DZ114 (5'-CCTTCTCA GGCCGGGTCCAGAAAGGATC-3') were oppositely directed to sequences flanking the *BglII* cloning site of pVL1393 and contained additional *Bsu36I* restriction sites (italics) at their 5' ends for circularization. DZ114 also contained an internal *SanDI* restriction site (underlined) for direct cloning into *SanDI*-linearized SeMNPV-US1 DNA. The template for PCR was purified AcMNPV bacmid bMON14272 (30) DNA from the Bac-to-Bac Kit (Gibco BRL). The resulting 8.5-kb PCR product was cloned into the 3.5-kb pCR-XL-TOPO vector (Invitrogen), digested with *Bsu36I*, self-ligated, and cloned into electrocompetent DH10 $\beta$  *E. coli* cells. The bacmid cloning vector obtained was designated BAC-Bsu36I, and its identity was verified by restriction analysis.

**Direct cloning of SeMNPV-US1 as bacmid.** SeMNPV-US1 DNA for direct cloning was purified using alkaline treatment of polyhedra and by previously described methods (36). Two micrograms of viral SeMNPV-US1 DNA was linearized at the polyhedrin locus by digestion with 10 U of *SanDI* (Stratagene) for 16 h at 37°C. The restriction enzyme was heat inactivated for 15 min at 65°C. One microgram of bacmid cloning vector BAC-Bsu36I was digested with 10 U of *SanDI* in a total volume of 35  $\mu$ l for 3 h at 37°C. The 8.5-kb vector was dephosphorylated using 1 U of HK Thermolabile Phosphatase (Epicentre). The enzymes were heat inactivated for 15 min at 65°C prior to gel purification of the linearized cloning vector DNA with Glassmax (Gibco BRL). Ligation was performed for 16 h at 15°C with approximately 500 ng of linearized SeMNPV DNA and 25 ng of linearized vector DNA in a total volume of 20  $\mu$ l using 6 U of T4 DNA ligase (Promega). Electrocompetent *E. coli* DH10 $\beta$  cells (Gibco BRL) were transformed with 2  $\mu$ l of ligation mix at 1.8 kV using a Bio-Rad Gene

Pulser. The transformed cells were recovered in SOC medium (40) for 45 min at 37°C and spread on agar plates containing kanamycin. An SeMNPV bacmid with the correct restriction profile was selected from 111 putative SeMNPV bacmid clones and was designated SeBAC10.

**Deletion of SeMNPV non-*hr ori* by ET recombination in *E. coli*.** For deletion mutagenesis of the active essential domain of the non-*hr ori* of SeMNPV-US1 bacmid SeBAC10, 68- to 70-bp primers were designed with 50-bp 5' ends flanking the deletion target region on the SeMNPV genome. Forward primer DZ153 was 5'-CATTTACTCGAAAACTGTACACTTCGTCAAAATAAATGAC GCAATATTTTAAAGGGACCAATAACTG-3', with a viral flanking sequence from nt 83237 to 83286 according to the SeMNPV complete genome sequence (16). Reverse primer DZ154 was 5'-ATTTCAAAATTAGAAATCAAACCCCA ATTTGCCGGCAACGTTTAAATATTTTCTGTGCGACGGTTAC-3', with a viral flanking sequence from nt 83981 to 83932. The locus to be deleted, which is the essential domain of the SeMNPV non-*hr ori*, is defined by two *SspI* sites (11). These *SspI* restriction sites are included in the primers (underlined). The 3' ends of the primers anneal to the chloramphenicol gene of pBeloBAC11 (41, 48) from nt 735 to 1671. PCR on pBeloBAC11 was performed using the Expand long-template PCR system (Roche) according to the manufacturer's protocol, giving a product of 1,036 bp. The PCR product was purified using the High pure PCR purification kit (Roche), cut with *DpnI* to eliminate residual pBeloBAC11 template, phenol-chloroform extracted, and ethanol precipitated. Approximately 0.5  $\mu$ g of PCR product was used for transformation of electrocompetent *E. coli* DH10 $\beta$  containing both SeBAC10 and homologous recombination helper plasmid pBAD- $\alpha$ - $\beta$ .

DH10 $\beta$  cells containing SeBAC10 were heat shock transformed with pBAD- $\alpha$ - $\beta$  (34) and subsequently made electrocompetent according to the method of Muylers et al. (34). Briefly, 70 ml of Luria-Bertani medium was inoculated with 0.7 ml of an overnight culture. At an optical density at 600 nm of 0.1 to 0.15, ET protein expression from pBAD- $\alpha$ - $\beta$  was induced by the addition of 0.7 ml of 10% L-arabinose. The cells were harvested at an optical density at 600 nm of 0.3 to 0.4 and made electrocompetent by three subsequent washes with ice-cold 10% glycerol. The cells were transformed with the purified PCR product in 2-mm-diameter electroporation cuvettes (Eurogentec) using a Bio-Rad Gene Pulser (2.3 kV, 25  $\mu$ F, 200  $\Omega$ ). The cells were resuspended in 1 ml of Luria-Bertani medium and incubated for 1 h at 37°C and subsequently spread on agar plates containing kanamycin and chloramphenicol. The altered genotype of the recombinant bacmid, designated SeBAC10 $\Delta$ nonhr, was confirmed by *PstI* digestion and PCR. The genomic *PstI*-I fragment of SeBAC10 (7,017 bp) was anticipated to be 286 bp bigger in SeBAC10 $\Delta$ nonhr, giving a fragment of 7,303 bp (Fig. 4). PCR was performed with forward primer DZ127 and reverse primer DZ128 as previously described. The PCR product of 1,213 bp was cloned into pGEM-Teasy (Promega) and completely sequenced, revealing that recombination had occurred precisely at the anticipated locus via the 50 flanking nucleotides.

**Reconstitution of the SeMNPV polyhedrin gene by pFastBAC1 donor plasmid.** To reconstitute the polyhedrin gene in SeMNPV bacmids SeBAC10 and SeBAC10 $\Delta$ nonhr, donor plasmid pFB1Sepol was constructed. The pFastBac1 vector (Gibco BRL) was digested with *SnaBI* and *HindIII* to delete the AcMNPV polyhedrin promoter and the multiple cloning site. The SeMNPV polyhedrin gene with its own promoter and the first putative transcription termination signal (46) was amplified by the Expand long-template PCR system (Roche) using forward primer DZ138, 5'-CCCCGGGTATATACTAGACGCGATTAC-3' (nt 135475 to 135494), and reverse primer DZ139, 5'-CCAGCTTTGTAATA CTTACCTTTTGTG-3' (nt 757 to 776), containing *SmaI* and *HindIII* restriction sites (italics), respectively. The resulting 930-bp fragment was cloned into a pGEM-Teasy vector (Promega), sequenced, and subsequently cloned as a *SmaI/HindIII* fragment into the pFastBac1 vector to generate pFB1Sepol. The protocol from the Bac-to-Bac manual (Gibco BRL) was followed to transpose the SeMNPV polyhedrin gene from pFB1Sepol into the *attTn7* transposon integration site of SeMNPV bacmids SeBAC10 and SeBAC10 $\Delta$ nonhr to generate SeBAC10ph and SeBAC10ph $\Delta$ nonhr, respectively.

**Transfection of SeMNPV bacmids.** Se301 cells were seeded in a six-well tissue culture plate (Nunc) at a confluency of 25% ( $5 \times 10^5$  cells). Transfection was performed with approximately 1  $\mu$ g of SeBAC10ph or SeBAC10ph $\Delta$ nonhr DNA using 10  $\mu$ l of Cellfectin (Gibco BRL). As a positive control, 1  $\mu$ g of SeMNPV-US1 DNA was transfected as well. After 5 and 7 days, polyhedra were formed by the cells transfected with SeMNPV-US1 and the bacmids, respectively. BV-containing supernatant (defined as P1) and infected cells were harvested 14 days posttransfection (90% polyhedron-containing cells).

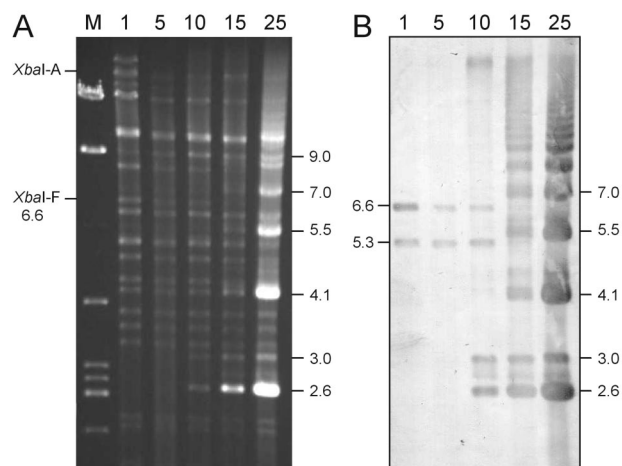


FIG. 1. Restriction profile of intracellular DNA of wild-type SeMNPV-US1 upon passing (P1 to 25) in Se301 insect cells. (A) DNA digested with *Xba*I and run in a 0.6% agarose gel. Passage numbers are indicated above the lanes, and the viral genomic *Xba*I-A and -F fragments are indicated on the left. Lane M contains a  $\lambda$ /EcoRI/*Bam*HI/*Hind*III DNA size marker. Sizes (in kilobases) of the hypermolar novel bands (2.6 to 7.0) and the novel 9-kb fragment are indicated on the right. (B) Southern blot using the SeMNPV non-*hr ori* (nt 83122 to 84048) as a probe. The viral genomic 6.6-kb *Xba*I-F (containing the non-*hr ori*) and an additional hybridizing 5.3-kb band are indicated on the left.

## RESULTS

**Serial passage of SeMNPV in Se301 insect cells.** SeMNPV-US1 was serially passed 25 times in the *S. exigua* cell line Se301 with BV from infectious hemolymph, defined as P0 inoculum. A decrease of polyhedron production was observed after fewer than five passages, indicating a dramatic passage effect. ICV DNA was purified and subjected to *Xba*I (Fig. 1A) digestion. A rapid reduction of the major genomic *Xba*I-A fragment was observed (Fig. 1A). At the same time, a novel *Xba*I fragment of about 9 kb became more abundant and was cloned and sequenced. This fragment was already present in the P1 DNA and appeared to consist of the remnants of the *Xba*I-A fragment as a result of a 26.5-kb deletion (from nt 15301 to 41759), according to the complete genome sequence of SeMNPV (16). The occurrence of mutants with deletions in this particular genomic region is a common phenomenon of SeMNPV infection in cell culture, but these deletions do not compromise BV or protein production (3, 10). In vivo, such deletion mutants also exist and can act as parasitic genotypes (33).

**Analysis of hypermolar bands.** Hypermolar fragments accumulated in Se301 cells from P10 onwards, and they were visualized as *Xba*I restriction fragments of 2.6 and 3.0 kb in agarose gels (Fig. 1A). From P15 onwards also, bands of 4.1, 5.5, and 7.0 kb and more became hypermolar. The abundant 2.6-, 3.0-, and 4.1-kb *Xba*I bands were cloned and sequenced, and it was found that the *Xba*I sites on either side of the cloned inserts corresponded to the SeMNPV *Xba*I restriction site at position 82132, according to the complete sequence of SeMNPV (16). Most interestingly, both the 2.6- and 3.0-kb fragments contained the entire SeMNPV non-*hr ori* origin of DNA replication (nt 83286 to 83932 [11]) and a junction of sequences flanking

this non-*hr ori* (Fig. 2). The borders and the junction of the 4.1-kb fragment appeared to be identical to those of the 2.6-kb fragment (Fig. 2B). The difference in size is a consequence of a duplicated non-*hr ori* present in this 4.1-kb fragment. Noteworthy is the presence of an overlapping stretch of 9 bp at the junction site in the 2.6- and 4.1-kb fragment (Fig. 2B), which in the complete SeMNPV genome is present on either side of the non-*hr ori*, leaving 2.6 kb in between. Because of the presence of a junction site and the same *Xba*I (position 82132) on either side of the fragments, it was concluded that the hypermolar fragments must exist in the ICV DNA preparation either as DNA minicircles or as tandem repeats in a larger concatenated form.

To investigate whether the other hypermolar bands of 5.5 kb, 7.0 kb, and more also contained the non-*hr ori*, a Southern blot was made with a non-*hr ori* probe. The result (Fig. 1B) showed that these fragments hybridized strongly to the probe, and therefore it was concluded that a range of molecules of different sizes containing the SeMNPV non-*hr ori* predominated upon serial passage.

In addition to the non-*hr ori* containing the genomic *Xba*I-F fragment of 6.6 kb, an unexpected additional band of 5.3 kb hybridizing to the non-*hr ori* probe (Fig. 1B [see also Fig. 3]) became submolar from P15 onwards. Sequencing revealed that the 5.3-kb fragment consisted of two joined, but distantly located, sequences from the SeMNPV genome (Fig. 2B). The ends of the fragment corresponded to *Xba*I sites at positions 82132 and 119846, respectively. The junction between the two fragments was formed by an overlapping sequence stretch of 19 bp, containing multiple GTC repeats, located at positions 86426 to 86446 and 118807 to 118780. The presence of this 5.3-kb band in the wild-type SeMNPV DNA was confirmed by Southern hybridization.

**Replicative-form hypermolar 2.6- and 3.0-kb *Xba*I fragments.** To investigate whether the abundant *Xba*I fragments of 2.6 and 3.0 kb exist as minicircles or as tandem repeats in a larger concatenated form, ICV DNA of P10 (at a stage when only the 2.6- and 3.0-kb bands were abundant) was subjected to partial digestion with *Xba*I, using increasing amounts of restriction endonuclease during digestion for 20 min. Hybridization was performed with the same non-*hr ori* probe as described above (Fig. 3). The partial *Xba*I digests of P10 viral DNA showed a stepladder of multimers of the 2.6- and 3.0-kb bands. This suggests that the accumulation of the SeMNPV non-*hr ori* occurs via high-molecular-weight concatemers of tandem repeats of different sizes. This is likely to be the case not only for the 2.6- and 3.0-kb fragments but also for the 4.1-, 5.5-, 7.0-kb, and larger fragments from P15 onwards.

**Serial passage of SeMNPV bacmids in Se301 insect cells.** A genetically homogeneous SeMNPV bacmid (SeBAC10) and a derived non-*hr ori* deletion mutant (SeBAC10 $\Delta$ nonhr) were constructed (Fig. 4) to determine whether non-*hr ori* concatemers are generated de novo in cell culture or preexist and become selectively amplified and whether virus stability might be enhanced by deletion of this non-*hr ori*.

Prior to serially passing the bacmid-derived BVs in cell culture, the polyhedrin gene was reintroduced. After transfection of Se301 cells, the BV-containing supernatant was defined as the P1 virus stock and was used to initiate serial undiluted passage. ICV DNA was purified and digested with



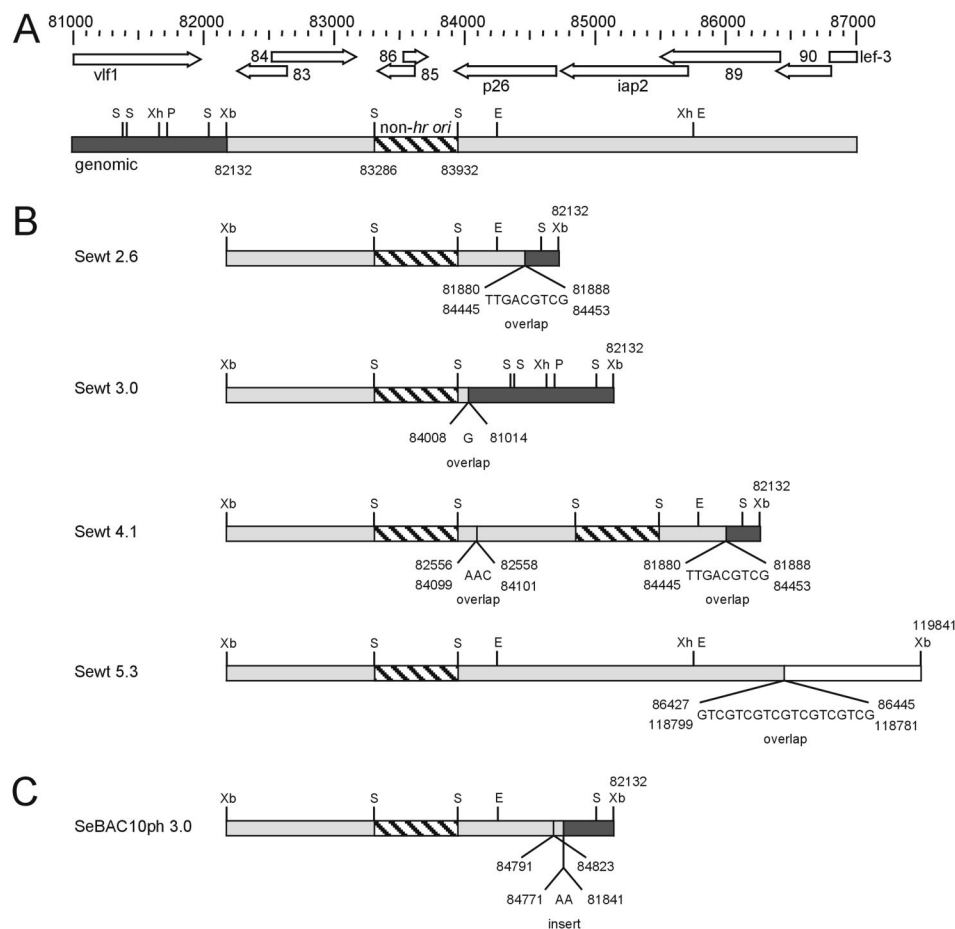


FIG. 2. Schematic overview of the genetic organization of hypermolar and other non-*hr ori* hybridizing bands compared to the complete SeMNPV genome. (A) Genetic organization of the genomic DNA with nucleotide positions according to the complete SeMNPV genome (16). Arrows represent the respective ORFs. Solid and light-grey boxes refer to sequences on either side of *Xba*I (Xb) 83132, containing *Ssp*I (S), *Pst*I (P), *Eco*RI (E), and *Xho*I (Xh) sites. The non-*hr ori* is presented as a hatched box between the two *Ssp*I sites (11). (B) Genetic arrangement of hypermolar 2.6-, 3.0-, and 4.1-kb fragments of SeMNPV-US1 (Sewt) and a nonhypermolar cohybridizing 5.3-kb fragment (genomic fragment of a SeMNPV deletion mutant) in the Southern blots. Nucleotide positions and sequence overlaps and insertions are indicated at the junction sites. (C) Genetic arrangement of the hypermolar 3.0-kb fragment of SeBAC10ph, containing two junctions.

*Xba*I. Similar to SeMNPV-US1 (this study and references 3 and 10), deletions in *Xba*I-A occurred for both bacmids SeBAC10ph and SeBAC10ph $\Delta$ nonhr (Fig 5). The deletion in SeBAC10ph $\Delta$ nonhr was mapped as a junction overlap of 3 nt (AAC) from 20162, 20163, or 20164 to 36396, 36397, or 36398, spanning open reading frames (ORFs) 17 to 35. From P6 onward, a small hypermolar *Xba*I fragment of 3.0 kb was visible in DNA preparations of SeBAC10ph (Fig. 5A). This fragment was cloned and sequenced and appeared to contain the non-*hr* origin of DNA replication and a junction sequence (Fig. 2c) also observed with the SeMNPV-US1 wild type (Fig. 2B). In contrast, the analysis of ICV DNA from SeBAC10ph $\Delta$ nonhr-infected cells did not reveal any accumulation of hypermolar fragments (Fig. 5B). SeBAC10ph $\Delta$ nonhr BV titers remained at higher levels throughout the entire period of serial passaging than those of SeBAC10ph and SeMNPV-US1 wild type (Fig. 6A). Polyhedron production of SeBAC10ph $\Delta$ nonhr remained constant for at least 20 passages, in contrast to SeBAC10ph (Fig. 6B). These results demonstrated that absence of the

non-*hr ori* strongly increased the stability of the SeMNPV genome in Se301 insect cells.

## DISCUSSION

The rapid accumulation of DI particles with DNA containing hypermolar non-*hr oris* appears to be an artifact of serial passage of SeMNPV in Se301 cells and the major cause of the decrease of virus and polyhedron production. Ultimately, these hypermolar molecules form the majority of the viral DNA. By partial digestion, we showed that these molecules exist as high-molecular-mass concatemers, in agreement with a supposed rolling-circle mechanism for baculovirus DNA replication (24, 28, 35, 50). Their rapid multiplication, together with previous data from in vitro replication assays (11), provides support for the view that the SeMNPV non-*hr ori* might be a genuine origin of DNA replication.

In order to elucidate whether these non-*hr ori* concatemers were newly formed in Se301 insect cells or, alternatively, only

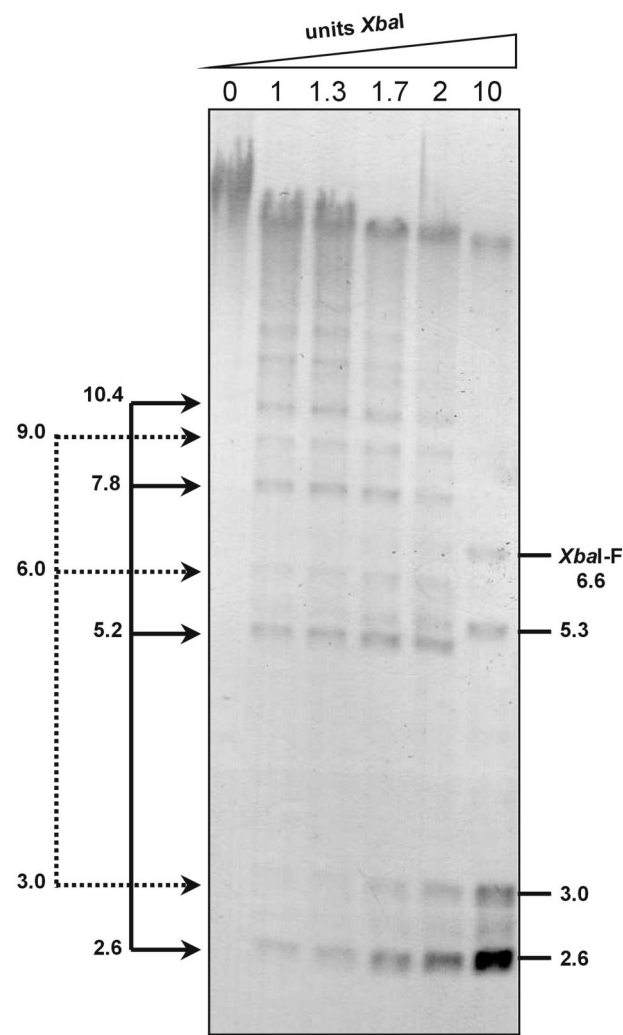


FIG. 3. Replicative form of the hypermolar 2.6- and 3.0-kb *Xba*I fragments by partial digestion of ICV SeMNPV-US1 DNA of P10, using increasing amounts of *Xba*I. On the right the genomic 6.6-kb *Xba*I-F and the additional 5.3-kb band as well as the hypermolar *Xba*I bands of 2.6 and 3.0 kb are indicated. On the left the multimers of the 2.6- and 3.0-kb *Xba*I fragments are indicated by arrows.

accumulated from a genetically heterogeneous wild-type isolate, a full-length infectious clone (bacmid) of SeMNPV was constructed. Such a bacmid could be stably maintained as a single-copy bacterial artificial chromosome in *E. coli* DH10 $\beta$  and used as a starting material in passage experiments in insect cells (30, 39). Transfection of Se301 insect cells with SeBAC10ph (with a reintroduced intact polyhedron gene) showed normal polyhedron production, but serial passage again resulted in a decrease of viral titers and polyhedron production and the rapid accumulation of non-*hr ori* containing molecules. Since the non-*hr ori* could be removed from the viral DNA without affecting virus replication, we must conclude that this non-*hr ori* is not essential. The two minor ORFs 85 and 86 within the non-*hr* are also nonessential. These ORFs do not have known baculovirus or other homologues, and it is unknown whether they are transcriptionally active. Furthermore, deletion of this non-*hr ori* strongly enhanced the genomic stability in cell culture. Transfection of Se301 cells with the non-*hr*

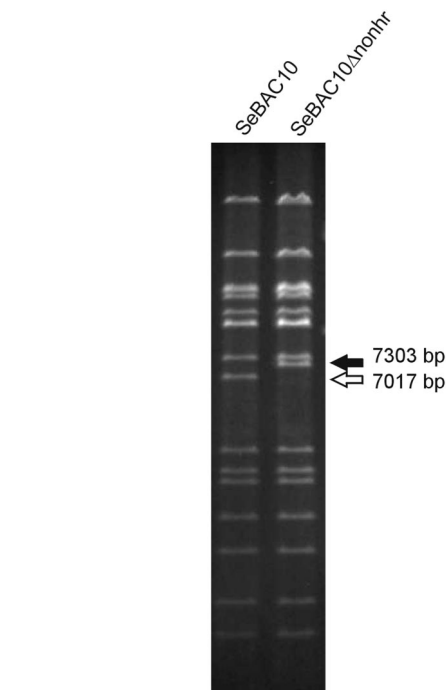


FIG. 4. Restriction profile (*Pst*I) of parental SeMNPV bacmid SeBAC10 and the non-*hr ori* deletion mutant SeBAC10 $\Delta$ nonhr. The genomic *Pst*I-I fragment containing the non-*hr ori* (7,017 bp) and the *Pst*I fragment with the *Cm*<sup>r</sup> gene insertion (7,303 bp) are indicated.

*ori* deletion mutant bacmid SeBAC10ph $\Delta$ nonhr gave normal polyhedron production and high viral titers (Fig. 6), which were maintained for up to at least 20 passages. Together with an unchanged restriction profile from P1 to P20, these results indicate that this recombinant has an enhanced stability in cell culture compared to both wild-type SeMNPV and the parental bacmid SeBAC10ph. The increased overall stability is probably

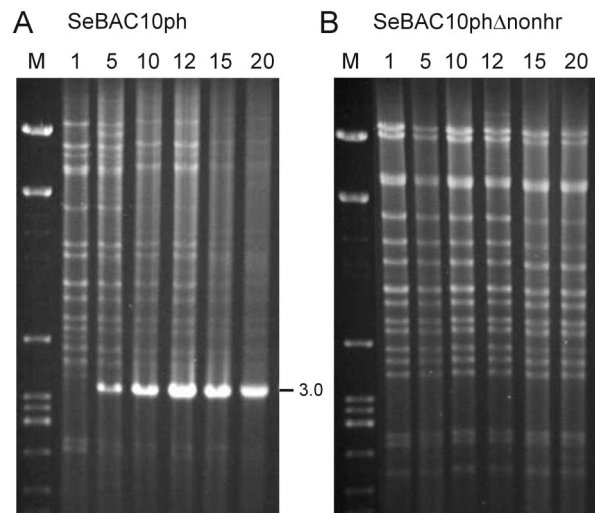


FIG. 5. Restriction profile (*Xba*I) of ICV DNA of serially passed SeMNPV bacmids SeBAC10ph (A) and SeBAC10ph $\Delta$ nonhr (B) in Se301 insect cells. Passage numbers (top) and the hypermolar band of 3.0 kb (SeBAC10ph) are indicated.

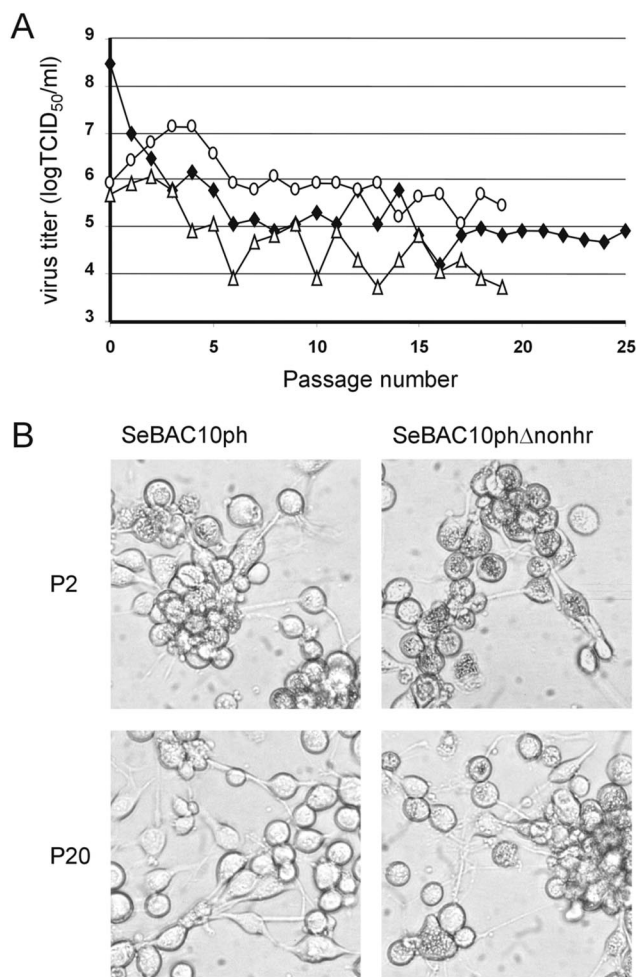


FIG. 6. (A) Titers of serially passaged BV of SeMNPV-US1 (◆), SeBAC10ph (△), and SeBAC10phΔnonhr (○). TCID<sub>50</sub>, 50% tissue culture infective dose. (B) Pictures of infected Se301 insect cells with SeBAC10ph and SeBAC10phΔnonhr at P2 and P20, respectively.

not due to an increase in intrinsic stability of the SeMNPV genome by itself, but rather a consequence of the lack of a *cis*-acting element (*non-hr ori*) that has a strong replicative advantage during baculovirus DNA replication.

Our findings with SeMNPV are in line with the results from Lee and Krell (26, 27), who demonstrated that AcMNPV DIs at P81 largely consisted of reiterations of about 2.8 kb of the standard genome, which was later shown to contain an active *non-hr* origin of DNA replication (7, 23). In addition, previous work on AcMNPV showed that DIs could also be enriched in *hrs* (21). These observations suggest that reiteration and predominance of baculovirus *hrs* but particularly *non-hr ori*s, which are complex structures comprising multiple direct and inverted repeats, are more common phenomena upon multiple passage and may contribute to a rapid passage effect. Single copies of *non-hr ori*s, which resemble eukaryotic *ori*s based on structural similarities (4), have been identified in many other baculovirus genomes by transient-replication assays (*Orgyia pseudotsugata* NPV [38] and *Spodoptera littoralis* MNPV [14, 15]) or based on sequence and structural similarity only (*Bom-*

*byx mori* NPV [24], *Busura suppressaria* NPV [13], and *Cydia pomonella* granulovirus [31]). The conservation of *non-hr ori* in baculovirus genomes implies an important (biological) role in virus replication and may be related to viral latency and persistence in insect populations. The resemblance to eukaryotic *ori*s suggests that baculoviruses may have obtained these sequences from the host genomes to be able to replicate in the insect without the requirement for virus-encoded replication factors.

At the junctions of *non-hr ori* concatenated molecules and junctions of major genomic deletions, sequence overlaps of 1 up to 19 bp were found, potentially involved in the causative recombination mechanism. The sequence of the additional 5.3-kb fragment in the Southern blots showed a 19-bp overlap at the deletion junction, consisting of multiple GTC repeats (Fig. 2B). These GTC repeats of up to 27 bp were found scattered throughout the SeMNPV genome on both strands and were more frequent than expected on a random basis. For the 2.6-kb *Xba*I fragment concatemers, a 9-bp TTGACGTCG overlap from flanking sequences was found at the junction site (Fig. 2B). Also, this repeat was found more frequently (12 times) in the genome than expected on a random basis (<1). The 9-bp overlap implies that the concatemers of the 2.6-kb fragment were generated during serial passage by looping and subsequent excision (homologous recombination) of *non-hr ori* containing genomic DNA, followed by continued replication and consequent concatenation of this intervening region. The same 9-bp overlap was found at the junction of the 4.1-kb fragment, which contains a duplicated *non-hr ori* (Fig. 2). This suggests that the 4.1-kb fragment was generated from the 2.6-kb molecule by another recombination event and rapidly became hypermolar because of the presence of two copies of the *non-hr ori*. This hypothesis is consolidated by the appearance of the 4.1-kb fragment in passages later than the 2.6-kb fragment (Fig. 1A). For the other junctions, smaller overlaps (1 or 3 bp) or even insertions were found, which was also demonstrated for AcMNPV DIs in an earlier study (39), suggesting that the same recombination mechanisms are involved.

The strategy of deletion of sequences that have a replicative advantage, accumulate upon serial passage, and interfere with virus and protein production (e.g., *non-hr ori*s) may now be applied for other baculoviruses as well and will contribute to the solution of problems associated with large-scale applications using baculovirus expression vectors for protein production in insect cells. In further studies we plan to map the sequences in the SeMNPV *non-hr ori* involved in the generation of DIs in more detail by reintroducing mutant SeMNPV *non-hr ori*s in SeMNPV bacmids. In addition, we want to investigate whether the deletion of the *non-hr ori* affects baculovirus persistence in vivo.

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